ENVIRONMENTAL CONTAMINANTS ENCYCLOPEDIA FUEL OIL NUMBER 5 ENTRY

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Like a library or most large databases (such as EPA's national STORET water quality database), this document contains information of variable quality from very diverse sources. In compiling this document, mistakes were found in peer reviewed journal articles, as well as in databases with relatively elaborate quality control mechanisms [366,649,940]. A few of these were caught and marked with a "[sic]" notation, but undoubtedly others slipped through. The [sic] notation was inserted by the editors to indicate information or spelling that seemed wrong or misleading, but which was nevertheless cited verbatim rather than arbitrarily changing what the author said.

Most likely additional transcription errors and typos have been added in some of our efforts. Furthermore, with such complex subject matter, it is not always easy to determine what is correct and what is incorrect, especially with the "experts" often disagreeing. It is not uncommon in scientific research for two different researchers to come up with different results which lead them to different conclusions. In compiling the Encyclopedia, the editors did not try to resolve such conflicts, but rather simply reported it all.

It should be kept in mind that data comparability is a major problem in environmental toxicology since laboratory and field methods are constantly changing and since there are so many different "standard methods" published by EPA, other federal agencies, state agencies, and various private groups. What some laboratory and field investigators actually do for standard operating practice is often a unique combination of various standard protocols and impromptu "improvements." In fact, the interagency task force on water methods concluded that [1014]:

It is the exception rather than the rule that water-quality monitoring data from different programs or time periods can be compared on a scientifically sound basis, and that...

No nationally accepted standard definitions exist for water quality parameters. The different organizations may collect data using identical or standard methods, but identify them by different names, or use the same names for data collected by different methods [1014].

Differences in field and laboratory methods are also major issues related to (the lack of) data comparability from media other than water: soil, sediments, tissues, and air.

In spite of numerous problems and complexities, knowledge is often power in decisions related to chemical contamination. It is therefore often helpful to be aware of a broad universe of conflicting results or conflicting expert opinions rather than having a portion of this information arbitrarily censored by someone else. Frequently one wants to know of the existence of information, even if one later decides not to use it for a particular application. Many would like to see a high percentage of the information available and decide for themselves what to throw out, partly because they don't want to seem uniformed or be caught by surprise by potentially important information. They are in a better position if they can say: "I knew about that data, assessed it based on the following quality assurance criteria, and decided not to use it for application." This is especially true for users near the end of long decision processes, such as hazardous site cleanups, lengthy ecological risk assessments, or complex natural resource damage assessments.

For some categories, the editors found no information and inserted the phrase "no information found." This does not necessarily mean that no information exists; it

simply means that during our efforts, the editors found none. For many topics, there is probably information "out there" that is not in the Encyclopedia. The more time that passes without encyclopedia updates (none are planned at the moment), the more true this statement will become. Still, the Encyclopedia is unique in that it contains broad ecotoxicology information from more sources than many other reference documents. No updates of this document are currently planned. However, it is hoped that most of the information in the encyclopedia will be useful for some time to come even with out updates, just as one can still find information in the 1972 EPA Blue Book [12] that does not seem well summarized anywhere else.

Although the editors of this document have done their best in the limited time available to insure accuracy of quotes or summaries as being "what the original author said," the proposed interagency funding of a bigger project with more elaborate peer review and quality control steps never materialized.

The bottom line: The editors hope users find this document useful, but don't expect or depend on perfection herein. Neither the U.S. Government nor the National Park Service make any claims that this document is free of mistakes.

The following is one chemical topic entry (one file among 118). Before utilizing this entry, the reader is strongly encouraged to read the README file (in this subdirectory) for an introduction, an explanation of how to use this document in general, an explanation of how to search for power key section headings, an explanation of the organization of each entry, an information quality discussion, a discussion of copyright issues, and a listing of other entries (other topics) covered.

See the separate file entitled REFERENC for the identity of numbered references in brackets.

HOW TO CITE THIS DOCUMENT: As mentioned above, for critical applications it is better to obtain and cite the original publication after first verifying various data quality assurance concerns. For more routine applications, this document may be cited as:

Irwin, R.J., M. VanMouwerik, L. Stevens, M.D. Seese, and W. Basham. 1997. Environmental Contaminants Encyclopedia. National Park Service, Water Resources Division, Fort Collins, Colorado. Distributed within the Federal Government as an Electronic Document (Projected public availability on the internet or NTIS: 1998).

<u>Fuel Oil Number 5 (No. 5 Fuel Oil, Bunker B Fuel Oil, Fuel Oil No.</u> 5, Navy Special Fuel Oil, NSFO, Residual Oil No. 5)

Brief Introduction:

Br.Class: General Introduction and Classification Information:

Fuel oils are comprised of mixtures of petroleum distillate hydrocarbons [363,499]. Fuel oil no. 5 consists of straight-run and cracked distillates and residuals, and contains aliphatics and aromatics [606]. ASTM specifications list two grades of No 5, one of which is lighter and under some climatic conditions may be handled and burned without preheating [498]. No. 5 fuel oil has two general categories: light and heavy [641]:

Light no. 5: This residual oil of intermediate viscosity is used in burners capable of handling fuel more viscous than fuel oil no. 4 without preheating. In some cases, preheating may be necessary in some types of equipment and in colder climates for handling.

Heavy no. 5: This residual fuel oil is more viscous than grade no. 5 (light), but is intended for similar use (see above). Preheating to 170-220 F is recommended before handling or use.

The viscous residuum fuel oils, numbers 5 and 6, sometimes referred to as bunker B and bunker C fuels, respectively, usually must be preheated before being burned [498].

Fuel oil numbers 4, 5, and 6 are commonly known as "residual oils" since they are manufactured in whole or in part from distillation residues from refinery processing [747]. These three heavy fuel oils are also known as gas oils and are composed of hydrocarbons ranging from C19 to C25 [872]. Residual oils are complex mixtures of relatively high molecular weight compounds and are difficult to characterize in detail. Molecular types include asphaltenes, polar aromatics, naphthalene aromatics, aromatics, saturated hydrocarbons and heteromolecules containing sulfur, oxygen, nitrogen, and metals [747].

Br. Haz: General Hazard/Toxicity Summary:

Many of the hazards to fish and wildlife from Fuel oil 5 are a result of the presence of polycyclic aromatic hydrocarbons (PAHs). Due to their relative persistence

and potential for various chronic effects (like carcinogenicity), PAHs, (and particularly the alkyl PAHs) can contribute to long term (chronic) hazards of fuel 5 in contaminated soils, sediments, groundwater, biological tissues (see PAHs entry). A NOAA protocol [828] GC/MS/SIM expanded scan for PAHs in a sample of fresh NSFO (Fuel Oil 5) revealed the presence of all 39 PAHs analyzed, with the PAHs and alkyl concentration being 0.6 ppm for Benzo(k)fluoranthene and the highest concentration being 4865 ppm for C2-Naphthalenes. A groundwater sample contaminated with weathered versions of the same NSFO product from Colonial National Historical Park also revealed the presence of all 39 PAHs and alkyl PAHs analyzed, with the lowest concentration being 39.7 ppt for Benzo(k)fluoranthene and the highest concentration being 48336 ppt for C4-Naphthalenes. In this Park Service groundwater investigation at Colonial National Historical Park, 92.4% of the total concentration values of the PAHs detected in groundwater were alkyl PAHs; parent compound PAHs analyzed in typical EPA scans accounted for only 7.6% of the PAHs and were below typical EPA scan detection limits, so EPA scans would have given false negatives. (see Chem.Detail section below for more details). provides a good example of why the standard EPA scans are inappropriate for hazard, damage, or risk assessment. Total naphthalenes in these groundwater samples exceeded 107,000 ppt (Roy Irwin, National Park Service, Personal Communication, 1996).

Regarding potential effects on humans, chronic effects of some of the constituents in fuel 5 (including PAHs such as naphthalenes) include changes in the liver and kidney [766]. Fuel oil #5 would be expected to be a skin, eye and respiratory irritant and a CNS depressant from inhalation of large amounts of the vapor or mist [606]. Prolonged or repeated contact with the skin may produce a defatting dermatitis with dryness and cracking [606].

One of the hazardous groups of compounds in fuel oils is PAHs (see PAHs entry). PAHs may be translocated in plants and may accumulate in plants grown in contaminated soil [40]. Presumably this also occurs in sediments and aquatic plants and therefore might impact herbivorous species of fish and wildlife. Although some research seems to indicate that interior portions of above-ground vegetables do not accumulate high concentrations of PAHs, plants do translocate PAHs from roots to other plant parts, such as developing shoots [40]. Some plants can evidently catabolize benzo(a)pyrene, but metabolic pathways have not been clearly defined. This is an important factor since when PAHs do degrade through metabolism, they often break down into even more toxic, carcinogenic, and mutagenic compounds [40]. Metabolic

transformations of PAHs into even more hazardous chemicals could also happen through microbial degradation of PAHs in soils or sediments. This provides an additional example of a situation where human health based standards are not protective of fish and wildlife, since it casts doubt on the environmental safety margin provided by EPA's human health-based soil guideline of =<100 ppm carcinogenic PAHs.

However, in a series of soil and hydrocultures of the higher plants, tobacco, rye, and radish, as well as algae cultures of lower plants (Chlorella Scenedesmus obligurus, vulgaris, Ankistrodesmus) /results indicate/ that certain polycyclic aromatic hydrocarbons (PAHs) growth-promoting effects on plants. Further, the degree of the promoting effect corresponded to the oncogenic activity of the hydrocarbon. The six polycyclic aromatic hydrocarbons found in plants were tested one at a time or in combination. Considerable growth-promotion was noted (near to 100% in some cases) with the effectiveness of hydrocarbons ranked as follows: (1) Benzo(a)pyrene (2) Benzo(a)anthracene (3) Indeno (1,2,3-cd)pyrene, Benzo(b)fluoranthene (4)Fluoranthene Benzo(ghi)perylene. [Graf W, Nowak W; Arch Hyg Bakt 150: 513-28 (1968) as cited in Health & Welfare Canada; Polycyclic Aromatic Hydrocarbons p.67 (1979) Report No. 80-EHD-50] [366].

See also: PAHs as a Group entry.

Br.Car: Brief Summary of Carcinogenicity/Cancer Information:

There is sufficient evidence for the carcinogenicity in experimental animals of residual (heavy) fuel oils and cracked residues derived from the oil refining of crude oil [747]. Residual (heavy) fuel oils are possibly carcinogenic to humans [747].

No studies were found for fuel oil #5 [606]. However, residual fuel oils contain appreciable concentrations of polynuclear aromatic hydrocarbons (PAHs) [747]. Fuel Oil Number 5 is typically about 75-80% Fuel Oil no. 6 [641], and the content of PAHs in Fuel Oil No. 6 is better documented (see: Fuel Oil No. 6). Dimethyl and trimethyl naphthalenes are important components of Fuel No. 6 (and also no. 5). Most blending stocks of residual fuel oils are likely to contain 5% or more of four- to six-ring condensed aromatic hydrocarbons [747]. This is important since some of the heavier PAHs are carcinogenic. When certain PAHs degrade through metabolism, they often break down into even more toxic, carcinogenic, and mutagenic

compounds [40]. Since more information is available on Fuel Oil 6, see also Fuel Oil Number 6 entry.

The debates on which PAHs, alkyl PAHs, and other aromatics found in this product to classify carcinogens, and the details of exactly how to perform both ecological and human risk assessments on the complex mixtures of PAHs typically found at contaminated sites, are likely to continue. There are some clearly wrong ways to go about it, but defining clearly right ways is more difficult. PAHs usually occur in complex mixtures rather than alone. Perhaps the most unambiguous thing that can be said about complex PAH mixtures is that such mixtures are often carcinogenic and possibly phototoxic. One way to approach site specific risk assessments would be to collect the complex mixture of PAHs and other lipophilic contaminants in a semipermeable membrane device (SPMD, also known as a fat baq) [894,895,896], retrieve the contaminant mixture from the SPMD, then test mixture for carcinogenicity, toxicity, phototoxicity (James Huckins, National Biological Service, and Roy Irwin, National Park Service, personal communication, 1996).

See also: PAHs as a Group entry.

Br.Dev: Brief Summary of Developmental, Reproductive, Endocrine, and Genotoxicity Information:

Some of the PAHs found in fuel oil are either AHH active or endocrine disruptors [561].

The results are mixed, but some reproductive and fetotoxic effects have been associated with a few of the compounds found in fuel no. 5 [766] (see entries on individual compounds for more details).

See Chem. Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture.

See also: PAHs as a group entry.

Br.Fate: Brief Summary of Key Bioconcentration, Fate, Transport, Persistence, Pathway, and Chemical/Physical Information:

Fuel Oil Number 5 is typically about 75-80% Fuel Oil no. 6 [641]. For concentrations of PAHs in the Fuel Oil 6, see Fuel Oil Number 6 entry. Dimethyl and trimethyl naphthalenes are important components of Fuel Oils Numbers 5 and 6.

Appreciable concentrations of polynuclear aromatic hydrocarbons (PAHs) are present in residual fuels because of the common practice of using both uncracked and cracked residues in their manufacture [747]. Most blending stocks of residual fuel oils are likely to contain 5% or more of four- to six-ring condensed aromatic hydrocarbons. The exact identities and concentrations of various PAHs depend on the nature and amount of the low viscosity blending stocks and the proportions of virgin and cracked residues [747].

Oil-soluble compounds of calcium, cerium, iron, or manganese may be added to residual fuel oils to improve combustion [747]. Concentrations vary with fuel oil, but typically range between 50 to 300 ppm weight of the active material ingredient [747].

Fuel oil number 5 is a heavy oil with little or no evaporation or dissolution potential (however, light fuel no. 5 would be more likely to experience some evaporation) [777]. As such, fuel oil 5 may be highly persistent, with the potential for long-term sediment contamination. In general, residual fuels are less acutely toxic relative to other oil types. They may weather slowly, and potentially sink depending on product density and water density (for a more complete discussion of sinking oil, see the Oil Spills entry). Impacts to waterfowl and fur-bearing animals can be severe during a spill of residual oil due to coating and ingestion [777].

Petroleum distillates in order of decreasing volatility include [363]:

- 1. Petroleum ether or benzene
- 2. Gasoline
- 3. Naphtha
- 4. Mineral spirits
- 5. Kerosene
- 6. Fuel oils
- 7. Lubricating oils
- 8. Paraffin wax
- 9. Asphalt or tar.

Synonyms/Substance Identification:

Residual Fuel Oil [560]
Navy Special [560]
Bunker B Fuel Oil [560]
Light Fuel Oil No. 5 [560]
Heavy Fuel Oil No. 5 [560]
FUEL OIL NO. 5 [606]
FUEL OIL #5 [606]
HEATING OIL NO. 5 [606]

HEATING OIL #5 [606] NO. 5 FUEL OIL [606] NO. 5 HEATING OIL [606] #5 FUEL OIL [606] #5 HEATING OIL [606] Fuel Oil UNSP [962]

Navy Special Fuel Oil (NSFO), Heavy Fuel Oil [ARCO MSDS Sheet] Commercial 535, ASTM No. 5 [ARCO MSDS Sheet]

Associated Chemicals or Topics (Includes Transformation Products):

See also individual entries:

Fuel Oil, General Fuel Oil Number 6 Oil Spills Petroleum, General PAHs as a group Naphthalene C1-Naphthalene C2-Naphthalene C3-Naphthalene C4-Naphthalene Biphenyl Acenaphthylene Acenaphthene Fluorene C1-Fluorene C2-Fluorene C3-Fluorene Anthracene Phenanthrene C1-Phenanthrene/anthracene C2-Phenanthrene/anthracene C3-Phenanthrene/anthracene C4-Phenanthrene/anthracene Dibenzothiophene C1-Dibenzothiophene C2-Dibenzothiophene C3-Dibenzothiophene Fluoranthene Pyrene C1-Fluoranthene/pyrene Benzo(a)anthracene Chrysene C1-Chrysene C2-Chrysene C3-Chrysene C4-Chrysene

Benzo(b)fluoranthene Benzo(k)fluoranthene Benzo(e)pyrene
Benzo(a)pyrene
Perylene
Indeno(1,2,3-c,d)pyrene
Dibenz(a,h)anthracene
Benzo(q,h,i)perylene

Site Assessment-Related Information Provided by Shineldecker (Potential Site-Specific Contaminants that May be Associated with a Property Based on Current or Historical Use of the Property) [490]:

Raw Materials, Intermediate Products, Final Products, and Waste Products Generated During Manufacture and Use:

- Benzene
- Creosote
- Ethyl benzene
- Polynuclear aromatic hydrocarbons
- Toluene
- Xylenes

Water Data Interpretation, Concentrations and Toxicity (All Water Data Subsections Start with "W."):

W.Low (Water Concentrations Considered Low):

No information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture.

W.High (Water Concentrations Considered High):

No information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture.

W.Typical (Water Concentrations Considered Typical):

No information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture.

W.Concern Levels, Water Quality Criteria, LC50 Values, Water Quality Standards, Screening Levels, Dose/Response Data, and Other Water Benchmarks:

W.General (General Water Quality Standards, Criteria, and Benchmarks Related to Protection of Aquatic Biota in General; Includes Water Concentrations Versus Mixed or

General Aquatic Biota):

No information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture.

W.Plants (Water Concentrations vs. Plants):

No information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture.

W.Invertebrates (Water Concentrations vs. Invertebrates):

No information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture.

W.Fish (Water Concentrations vs. Fish):

No information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture.

W.Wildlife (Water Concentrations vs. Wildlife or Domestic
Animals):

No information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture.

W.Human (Drinking Water and Other Human Concern Levels):

No information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture.

W.Misc. (Other Non-concentration Water Information):

No information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture.

Sediment Data Interpretation, Concentrations and Toxicity (All Sediment Data Subsections Start with "Sed."):

Sed.Low (Sediment Concentrations Considered Low):

No information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture.

Sed.High (Sediment Concentrations Considered High):

No information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture.

Sed.Typical (Sediment Concentrations Considered Typical):

No information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture.

Sed.Concern Levels, Sediment Quality Criteria, LC50 Values, Sediment Quality Standards, Screening Levels, Dose/Response Data and Other Sediment Benchmarks:

Sed.General (General Sediment Quality Standards, Criteria, and Benchmarks Related to Protection of Aquatic Biota in General; Includes Sediment Concentrations Versus Mixed or General Aquatic Biota):

No information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture.

Sed.Plants (Sediment Concentrations vs. Plants):

No information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture.

Sed.Invertebrates (Sediment Concentrations vs. Invertebrates):

No information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture.

Sed.Fish (Sediment Concentrations vs. Fish):

No information found; see Chem.Detail section for compounds in this product, then see individual

compound entries for summaries of information on individual components of this mixture.

Sed.Wildlife (Sediment Concentrations vs. Wildlife or Domestic Animals):

No information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture.

Sed. Human (Sediment Concentrations vs. Human):

No information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture.

Sed.Misc. (Other Non-concentration Sediment Information):

No information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture.

Soil Data Interpretation, Concentrations and Toxicity (All Soil Data Subsections Start with "Soil."):

Soil.Low (Soil Concentrations Considered Low):

No information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture.

Soil.High (Soil Concentrations Considered High):

Seven large and medium size west coast ports were surveyed during August 1990 to determine their involvement with hydrocarbon contaminated soils and activities associated with the characterization and remediation of these soils [735]. All ports surveyed indicated that they have hydrocarbon contaminated soil problems [735]. At one site, a soil investigation revealed one or more of four underground petroleum pipelines, all idle or abandoned, near the center of the redevelopment area may have leaked. The presence of petroleum contamination in the soil was confirmed. The petroleum could not be identified, but appeared to be of a heavy petroleum type (diesel, bunker oil (bunker B is fuel oil no. 5, bunker C is fuel oil no. 6), or possibly very weathered crude) rather than gasoline [735]:

Total Petroleum Hydrocarbon (TPH)	
EPA Method 418.1	69,300
Total Petroleum Hydrocarbon (TPH)	
EPA Method 8015 modified for diesel	43,000
Benzene	40.7
Toluene	102
Xylene	67
Ethylbenzene	171

Soil.Typical (Soil Concentrations Considered Typical):

No information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture.

Soil.Concern Levels, Soil Quality Criteria, LC50 Values, Soil Quality Standards, Screening Levels, Dose/Response Data and Other Soil Benchmarks:

Soil.General (General Soil Quality Standards, Criteria, and Benchmarks Related to Protection of Soil-dwelling Biota in General; Includes Soil Concentrations Versus Mixed or General Soil-dwelling Biota):

Soil.Plants (Soil Concentrations vs. Plants):

No information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture. See also: Br.Haz section above.

Soil.Invertebrates (Soil Concentrations vs. Invertebrates):

No information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture.

Soil.Wildlife (Soil Concentrations vs. Wildlife or Domestic Animals):

Acute Toxicity Highlights from RTECS [365]:

LDLO/LCLO - LOWEST PUBLISHED LETHAL DOSE/CONC RABBIT: LDLo; ROUTE: Skin; DOSE: 5200 mg/kg; REFERENCE: Acute Toxicity Data. Journal of the American College of Toxicology, Part B 1:139, 1990.

IRRITATION SKIN - STANDARD DRAIZE TEST, RABBIT, ROUTE: Skin; DOSE: 500 mg; REACTION: mild; REFERENCE: Acute Toxicity Data. Journal of the American College of Toxicology, Part B 1:139, 1990.

Soil. Human (Soil Concentrations vs. Human):

No information found on this complex and variable mixture. See Chem.Detail section for chemicals found in this product, then look up information on each hazardous compound. Some individual compounds found in petroleum products have low-concentration human health benchmarks for soil (see individual entries).

Soil.Misc. (Other Non-concentration Soil Information):

As of 1996, several States were considering allowing natural attenuation (the "do nothing and let nature clean up the mess through bioremediation" option) to proceed near leaking storage tanks in situations where drinking water was not being impacted and where human rather than environmental resources were the main resources in the immediate area (Roy Irwin, National Park Service, personal communication, 1996).

Others would point out that fuel oil spills into soils are not necessarily a trivial environmental threat related to ecotoxicology (emphasis on living things other than humans), due to the many hazardous compounds in this product (see Chem.Detail section below).

No other information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture.

Tissue and Food Concentrations (All Tissue Data Interpretation Subsections Start with "Tis."):

Tis.Plants:

A) As Food: Concentrations or Doses of Concern to Living Things Which Eat Plants:

No information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture.

B) Body Burden Residues in Plants: Typical, Elevated, or of Concern Related to the Well-being of the Organism

Itself:

No information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture.

Tis.Invertebrates:

A) As Food: Concentrations or Doses of Concern to Living Things Which Eat Invertebrates:

No information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture.

B) Concentrations or Doses of Concern in Food Items Eaten by Invertebrates:

No information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture.

C) Body Burden Residues in Invertebrates: Typical, Elevated, or of Concern Related to the Well-being of the Organism Itself:

No information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture.

Tis.Fish:

A) As Food: Concentrations or Doses of Concern to Living Things Which Eat Fish (Includes FDA Action Levels for Fish and Similar Benchmark Levels From Other Countries):

No information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture.

B) Concentrations or Doses of Concern in Food Items Eaten by Fish:

No information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture.

C) Body Burden Residues in Fish: Typical, Elevated, or of

Concern Related to the Well-being of the Organism Itself:

No information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture.

Tis.Wildlife: Terrestrial and Aquatic Wildlife, Domestic Animals and all Birds Whether Aquatic or not:

A) As Food: Concentrations or Doses of Concern to Living Things Which Eat Wildlife, Domestic Animals, or Birds:

No information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture.

B) Concentrations or Doses of Concern in Food Items Eaten by Wildlife, Birds, or Domestic Animals (Includes LD50 Values Which do not Fit Well into Other Categories, Includes Oral Doses Administered in Laboratory Experiments):

LD50/LC50 - LETHAL DOSE/CONC 50% KILL, RAT, LD50; ROUTE: Oral; DOSE: 5300 mg/kg; REFERENCE: Acute Toxicity Data. Journal of the American College of Toxicology, Part B 1:139, 1990 [365].

C) Body Burden Residues in Wildlife, Birds, or Domestic Animals: Typical, Elevated, or of Concern Related to the Well-being of the Organism Itself:

No information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture.

Tis.Human:

A) Typical Concentrations in Human Food Survey Items:

No information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture.

B) Concentrations or Doses of Concern in Food Items Eaten by Humans (Includes Allowable Tolerances in Human Food, FDA, State and Standards of Other Countries:

No information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on

individual components of this mixture.

C) Body Burden Residues in Humans: Typical, Elevated, or of Concern Related to the Well-being of Humans:

No information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture.

Tis.Misc. (Other Tissue Information):

No information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture.

Bio.Detail: Detailed Information on Bioconcentration, Biomagnification, or Bioavailability:

No information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture.

Interactions:

No information found; see Chem.Detail section for compounds in this product, then see individual compound entries for summaries of information on individual components of this mixture.

Uses/Sources:

The viscous residuum fuel oils, numbers 5 and 6, sometimes referred to as bunker fuels, are used in furnaces and boilers of utility power plants, ships, locomotives, metallurgical operations, and industrial power plants, and usually must be preheated before being burned [498].

Forms/Preparations/Formulations:

Fuel oil no. 5 can be prepared by combining 20 to 25 percent no. 2 fuel oil with 75 to 80 percent no. 6 fuel oil [641].

Chem.Detail: Detailed Information on Chemical/Physical Properties:

Caution: Every individual petroleum product has a unique "fingerprint," or distinct set of constituents most commonly identified by a gas chromatograph analysis. Due to the varying properties of the same general category of a petroleum product (each source and weathering stage of a No. 5 fuel oil

has a unique gas chromatograph "fingerprint"), careful assessment of the toxicity, specific gravity, and other physical characteristics of each individual oil must be taken into consideration to determine the exact effects of the product on the environment. Therefore, the below comments on No. 5 fuel oils are to be considered as representative, but not absolute values typical of every batch of the product with the same name.

Since PAHs are important hazardous components of this product, risk assessments should include analyses of PAHs and alkyl PAHs utilizing the NOAA protocol expanded scan [828] or other rigorous GC/MS/SIM methods.

The heavy fuel oils (fuels oils number 4,5, and 6) contain 15-40% aromatics, dominated by alkyl phenanthrenes and naphthalenes [872].

Fuel Oil Number 5 is typically about 75-80% Fuel Oil no. 6 [641]. For concentrations of PAHs in Fuel Oil no. 6, see Fuel Oil No. 6 entry. Dimethyl and trimethyl naphthalenes are important components of Fuel No. 6 (and also No. 5).

Physicochemical information from ORNL [875]:

Log KOW [875]: 3.3 to 7.06

Henry's Law Const [875]: 5.9E-05 to 7.4

Solubility in Water [875]: ~5 mg/L.

Odor [875]: like kerosene.

Vapor Pressure: 2.12 to 26.4 mm Hg at 21 degrees C [875].

SOLUBILITY [498]: Fuel oil is insoluble (sic, actually "relatively insoluble") in water.

NOTE on Solubility: No exact numbers can be given for solubilities of fuel oil in water because the composition of an oil varies from refinery to refinery. Generally, hydrocarbons of a lower molecular weight are more soluble than those of a higher molecular weight. Branching of hydrocarbon isomers, as well as ring formation, also tends to increase solubility. For two rings with the same carbon number, an unsaturated ring is more soluble in water than a saturated ring. The solubility of hydrocarbons in sea water is less than in fresh water. Also, an increase in temperature will greatly increase the amount of hydrocarbons which dissolve in water. Turbulence will also increase the rate of solubility [641].

API GRAVITY (60/60 degrees F) [560]:

NOTE: Created by the American Petroleum Institute (API), API gravity is an arbitrary scale expressing the gravity

or density of liquid petroleum products [637]. This scale was created in order to compare the densities of various oils. API gravity = (141.5/specific gravity [60/60 degrees F]) - 131.5, where specific gravity [60/60 degrees F] is the oil density at 60 degrees F divided by the density of water at 60 degrees F.

min. 11.5

METAL CONTENT [560]:

Other Metals (ppm):

Vanadium	152
Nickel	29.0
Chromium	1.076
Cobalt	0.198
Iron	24.0
Molybdenum	0.117
Manganese	0.248
Zinc	1.73
Copper	0.321

SPECIFIC GRAVITY [498]:

Less than 1 (Fuel oils Nos 1, 2, 4, 5).

DENSITY:

Density (g/mL) [560]:

For temperatures of oil (T) between 0 and 30 C: Density = 0.97871 - 0.000710 T

NOTE: The densities of crude oils and oil products are dependent on the temperature and degree of weathering (i.e., as temperature goes down and/or the fraction of weathered oil increases, density increases). The following density values are at "0% Weathering Volume" - in other words, fresh No. 5 fuel oil.

```
Temp( C) Density (at 0% Weathering Volume)
0 0.932 to 0.957
15 0.923 to 0.948
16 0.935
20 0.920 to 0.945
30 0.913 to 0.938
```

The Density is Less than 1 g/cm(3) for Fuel oils Nos 1, 2, 4, and 5 [498].

PAHs in NSFO (Fuel Oil 5):

A NOAA protocol [828] GC/MS/SIM expanded scan for PAHs in a sample of fresh NSFO (Fuel Oil 5) revealed the presence of all 39 PAHs and alkyl PAHs analyzed, with the lowest concentration being 0.6 ppm for Benzo(k)fluoranthene and the highest concentration being 4865 ppm for C2-Naphthalenes. groundwater sample contaminated with weathered versions of the same NSFO product from Colonial National Historical Park also revealed the presence of all 39 PAHs and alkyl PAHs analyzed, the lowest concentration being 39.7 ppt Benzo(k)fluoranthene and the highest concentration being 48336 ppt for C4-Naphthalenes (Chuck Rafkind, National Park Service, Personal Communication, 1996). Total naphthalenes in these groundwater samples exceeded 107000 ppt, far above the 2.8 ppt cancer risk (10-6 level) benchmark given for human health (see naphthalene section).

Details of PAH content (ng/mg or ppm) in one fresh sample of NSFO (Fuel Oil 5, Chuck Rafkind, National Park Service, Personal Communication, 1996):

```
Naphthalene:
                               34.3
C1-Naphthalene:
                               4086.9
                               4865.4
C2-Naphthalene:
                               4793.7
C3-Naphthalene:
C4-Naphthalene:
                               2688.5
Biphenyl:
                               3.5
Acenaphthylene:
                               4.1
Acenaphthene:
                               111.2
Fluorene:
                               216.0
C1-Fluorene:
                               658.8
C2-Fluorene:
                               1277.0
C3-Fluorene:
                               1243.8
Anthracene:
                               96.4
Phenanthrene:
                               778.2
C1-Phenanthrene/anthracene:
                               2116.3 (includes both)
C2-Phenanthrene/anthracene:
                               2716.7
C3-Phenanthrene/anthracene:
                               1923.3
C4-Phenanthrene/anthracene:
                               820.5
Dibenzothiophene:
                               25.7
C1-Dibenzothiophene:
                               1396.1
C2-Dibenzothiophene:
                               2155.9
C3-Dibenzothiophene:
                               1975.5
Fluoranthene:
                               31.6
                               177.9
Pyrene:
                               566.1
C1-Fluoranthene/pyrene:
Benzo(a)anthracene:
                               41.1
                               74.3
Chrysene:
C1-Chrysene:
                               312.1
                               370.8
C2-Chrysene:
                               29.9
C3-Chrysene:
C4-Chrysene:
                               19.7
Benzo(b)fluoranthene
                               11.0
Benzo(k)fluoranthene:
                               0.6
                               29.8
Benzo(e)pyrene:
```

Benzo(a)pyrene: 19.3
Perylene 10.6
Indeno(1,2,3-c,d)pyrene: 2.3
Dibenz(a,h)anthracene: 4.0
Benzo(g,h,i)perylene: 11.4

Note: The above PAHs and alkyl PAHs were analyzed by a GC/MS/SIM NOAA protocol [828] modified with methylene chloride extraction for use with water samples (Guy Denoux, Geochemical and Environmental Research Group, Texas A&M University, personal communication, 1996).

Details of PAH content (ng/L or ppt, compare to the above listed ppm concentrations by dividing the below-listed ppt concentrations below by 1,000,000) in one sample of groundwater subjected to long term contamination of NSFO (Fuel Oil 5), possibly mixed with some JP-4, motorgas, and JP-8, Colonial National Historical Park Groundwater Site MW-10 (Chuck Rafkind, National Park Service, Personal Communication, 1996):

```
530.8
Naphthalene:
C1-Naphthalene:
                              2463.7
C2-Naphthalene:
                             12044.7
C3-Naphthalene:
                             45345.1
C4-Naphthalene:
                             48336.8
Biphenyl:
                              129.7
                              81.2
Acenaphthylene:
Acenaphthene:
                              1517.6
                              1229.3
Fluorene:
C1-Fluorene:
                              11424.5
C2-Fluorene:
                              28680.7
C3-Fluorene:
                              32509.9
Anthracene:
                              1972.5
Phenanthrene:
                              7136.3
C1-Phenanthrene/anthracene:
                              31377.0 (includes both)
C2-Phenanthrene/anthracene:
                              49447.3
C3-Phenanthrene/anthracene:
                              41754.1
                              22250.2 "
C4-Phenanthrene/anthracene:
Dibenzothiophene:
                              8377.8
C1-Dibenzothiophene:
                              24742.0
C2-Dibenzothiophene:
                              44033.0
C3-Dibenzothiophene:
                              43900.3
Fluoranthene:
                              818.8
Pyrene:
                              5900.6
                              16248.3 (includes both)
C1-Fluoranthenes/pyrenes:
Benzo(a)anthracene:
                              1053.5
                              1817.1
Chrysene:
C1-Chrysene:
                              7398.8
C2-Chrysene:
                              9910.6
C3-Chrysene:
                             1048.5
                              625.9
C4-Chrysene:
Benzo(b)fluoranthene
                              399.2
```

39.7
1062.3
602.7
428.6
106.8
117.1
421.4

Note: The above PAHs and alkyl PAHs were analyzed by a GC/MS/SIM NOAA protocol [828] modified with methylene chloride extraction for use with water samples (Guy Denoux, Geochemical and Environmental Research Group, Texas A&M University, personal communication, 1996).

Additional Detailed Physicochemical information from Environment Canada [560]:

NOTE: In this section, for properties with more than one value, each value came from its own source; in other words, if API Gravity at 60 F was measured several times and several different answers were obtained, all of the answers are provided [560]:

VISCOSITY

NOTE: Viscosity is a measure of the internal friction or the resistivity to flow of a liquid [637]. The viscosities of crude oils and oil products are dependent on the temperature and degree of weathering (i.e., as temperature goes down and/or the fraction of weathered oil increases, viscosity increases). The following viscosity values are at "0% Weathering Volume" - in other words, fresh No. 5 fuel oil.

Kinematic Viscosity (mm2/sec or cSt):

Temp(C)	Light	Heavy
10	200 to 473	> 473
15	152 to 313	> 313
20	123 to 233	> 233
25	100 to 165	165 to 327
30	74 to 125	125 to 200
38	26.4 to 65	65 to 194
40	40 to 75	75 to 100

NOTE: Data obtained from a graph (Curl 77)

Pour Point (degrees C):

Pour point is the lowest temperature at which an oil sample is observed to flow when cooled under prescribed conditions. It is affected by weathering (i.e., the larger the fraction of oil weathered, the higher the pour point).

```
\max_{-17.8}
```

FIRE AND REACTIVITY

NOTE: Flash point is the lowest temperature at which vapors arising from the oil will ignite momentarily (i.e., flash) on application of a flame under specific conditions [637].

```
Flash Point (degrees C):
    > 54
    min 60
    min 55
    > 54.4

Flammability Limits (Volume %):
    in air: 1 to 5
```

DISTILLATION

NOTE: Distillation data provides an indication of an oil's volatility and relative component distribution. Distillation data is reported as volume % recovered.

Boiling Range (degrees C):

218 to > 570

SENSATION

Color Brown

OTHER

Reid method Vapor Pressure (kPa):

Temp(C) Pressure 37.8 0.689 (estimated)

Information from OHM/TADS [499]:

FIRE/EXPLOSION/CORROSION HAZARDS

Fire hazard:

Flammability: MODERATE. COMBUSTION WITH HEATING MODERATE

HAZARD

Standard codes: NFPA - -,2,0

Toxic combustion products: NO GREAT HAZARD

Flash point (degrees C): 66

Autoignition temperature (degrees C): 408

Fate.Detail: Detailed Information on Fate, Transport, Persistence, and/or Pathways:

See Br. Fate section above for additional information.

Persistency [499]: Loss of fuel oil after 40 hour in bubbler apparatus - 2.% Evaporated, .005% Dissolved.

Laboratory and/or Field Analyses:

As documented in sections of above, Fuel oil 5 contains a wide variety of PAHs. In fact, both fresh fuel oil 5 and a groundwater sample contaminated with weathered versions of the same Fuel Oil 5 revealed the presence of all 39 PAHs and alkyl PAHs analyzed by a GC/MS/SIM NOAA protocol [838] modified for use with water samples (see Chem.Detail section above).

Appreciable concentrations of PAHs are present in residual fuels because on the common practice of using both uncracked and cracked residues in their manufacture [747].

Many lab methods have been used to determine PAHs, BTEX compounds, and other common components of this fuel [861], but when potential biological effects are to being considered, many of the methods historically used have been determined to be inferior to the NOAA protocol expanded scan [828] being recommended by many risk assessment experts in 1996. Many historically used methods, including EPA standard semi-volatile scan number 8270, do not cover important alkyl PAHs and do not utilize low-enough detection limits to use in ecological risk assessments. See also: PAHs as a Group entry.

Recent (1991) studies have indicated that EPA approved methods used for oil spill assessments (including total petroleum hydrocarbons method 418.1, semivolatile priority pollutant organics methods 625 and 8270, and volatile organic priority pollutant methods 602, 1624, and 8240) are all inadequate for generating scientifically defensible information for Natural Resource Damage Assessments [468]. These general organic chemical methods are deficient in chemical selectivity (types of constituents analyzed) and sensitivity (detection limits); the deficiencies in these two areas lead to an inability to interpret the environmental significance of the data in a scientifically defensible manner [468].

If a Park Service groundwater investigation at Colonial National Historical Park performed in response to contamination by Fuel Oil 5 had utilized EPA semi-volatile scan 8270 or any of the other typical EPA scans (625, etc.) all of which only include parent compounds and typically utilize detection limits in the 170-600 ppb range, the false conclusion reached would have been that no PAHs were present in significant (detection limit) amounts. This false negative conclusion would have been made because the parent compound PAHs present constituted only 7.6% of the PAHs detected in groundwater by the expanded scan [828], and the highest concentration found for any parent compound was 8.4 ppb, far below the detection limits used on the older standard EPA scans.

Utilizing the NOAA protocol expanded scan [828], it was determined that 92.4% of the total concentration values of the PAHs detected in groundwater were alkyl PAHs, and that all 39 PAHs and alkyl PAHs were present. Of course, all 39 PAHs were also present in the fresh product, in much higher concentrations, and also having alkyl compounds with the highest percentage of higher values compared to parent compounds (see Chem.Detail section above for more details).

Due to the presence of light aromatics and PAHs in fuel oils, we recommend the following decision tree:

Decision Tree (dichotomous key) for selection of lab methods for measuring contamination from midrange to heavy crude oils, number 4 and heavier fuel oils, bunker C and all other oils considered to be heavy):

- 1a. Your main concern is biological effects of petroleum products......2
- 1b. Your main concern is cleanup or remediation but no ecological or human resources are at risk.............3
- 2b. The resource at risk is something else......5
- The spilled substance is a fresh* oil product of known 3a. composition: If required to do so by a regulatory authority, perform whichever Total Petroleum Hydrocarbon (TPH) analysis specified by the regulator. However, keep in mind that due to its numerous limitations, the use of the common EPA method 418.1 for Total Petroleum Hydrocarbons is not recommended as stand-alone method unless the results can first be consistently correlated (over time, as the oil ages) with the better NOAA protocol expanded scan*** for polycyclic aromatic hydrocarbons (PAHs) and alkyl PAHs. If not required to perform an EPA method 418.1-based analysis for TPH, instead perform a Gas Chromatography/Flame Ionization Detection (GC/FID) analysis for TPH using the spilled substance as a calibration standard. GC/FID methods can be sufficient for screening purposes when the oil contamination is fresh*, unweathered oil and when one is fairly sure of the source [657]. If diesel 1D was spilled, perform TPH-D (1D) using

California LUFT manual methods (typically a modified EPA method 8015) [465] or a locally available GC/FID method of equal utility for the product spilled. However, no matter which TPH method is used, whether based on various GC/FID or EPA method 418.1 protocols, the investigator should keep in mind that the effectiveness of the method typically changes as oil ages, that false positives or false negatives are possible, and that the better Gas Chromatography-Mass Spectrometry-Selected Ion Mode (GC/MS/SIM) scans (such as the NOAA expanded scan***) should probably be performed at the end of remediation to be sure that the contamination has truly been cleaned up.

- 3b. The spilled product is not fresh* or the contamination is of unknown or mixed composition......6
- 4. Analyze for Benzene, Toluene, Ethyl Benzene, and Toluene (BTEX) compounds in water as part of a broader scan of volatiles using EPA GC/MS method 8240. The standard EPA GC/MS method 8240 protocol will be sufficient for some applications, but the standard EPA method 8240 (and especially the less rigorous EPA BTEX methods such as method 8020 for soil and method 602 for water) are all inadequate for generating scientifically defensible information for Natural Resource Damage Assessments [468]. The standard EPA methods are also inadequate for risk assessment purposes. Thus, when collecting information for possible use in a Natural Resource Damage Assessment or risk assessment, it is best to ask the lab to analyze for BTEX compounds and other volatile oil compounds using a modified EPA GC/MS method 8240 method using the lowest possible Selected Ion Mode detection limits and increasing the analyte list to include as many alkyl BTEX compounds as possible. Also analyze surface or applicable) ground water samples for polycyclic aromatic hydrocarbons (PAHs) and alkyl PAHs using the NOAA protocol expanded scan*** modified for water samples using methylene chloride extraction. Ιf the contaminated water groundwater, before the groundwater is determined to be remediated, also analyze some contaminated sub-surface soils in contact with the groundwater for BTEX compounds (EPA GC/MS method 8240), and PAHs (NOAA protocol expanded scan***). The magnitude of any residual soil contamination will provide insight about the likelihood of recontamination of groundwater resources through equilibria partitioning mechanisms moving contamination from soil to water.
- 5a. The medium of concern is sediments or soils......6
- 5b. The medium of concern is biological tissues......7
- 6. Perform the NOAA protocol expanded scan*** for polycyclic aromatic hydrocarbons (PAHs) and alkyl PAHs. If there is any reason to suspect fresh* or continuing contamination of soils or sediments with lighter volatile compounds, also perform EPA

GC/MS method 8240 using the lowest possible Selected Ion Mode (SIM) detection limits and increasing the analyte list to include as many alkyl Benzene, Toluene, Ethyl Benzene, and Xylene (BTEX) compounds as possible.

- 7a. The problem is direct coating (oiling) of wildlife or plants with spilled oil product.....8
- 7b. The problem is something else.....9
- 8. Perform NOAA protocol expanded scan*** for polycyclic aromatic hydrocarbons (PAHs) and alkyl PAHs and/or GC/FID fingerprinting of the coating oil only if necessary to identify the source or exact oil. If the source is known and no confirmation lab studies are necessary: dispense with additional chemical laboratory analyses and instead document direct effects of coating: lethality, blinding, decreased reproduction from eggshell coating, etc., and begin cleaning activities if deemed potentially productive after consolations with the Fish and Wildlife Agencies.
- 9a. The concern is for impacts on water column organisms (such as fish or plankton)......10
- 9b. The concern is for something else (including benthic organisms)......11
- If exposure to fish is suspected, an HPLC/Fluorescence scan 10. for polycyclic aromatic hydrocarbon (PAH) metabolites in bile may be performed to confirm exposure [844]. For bottomdwelling fish such as flounders or catfish, also analyze the bottom sediments (see Step 6 above). Fish which spend most of their time free-swimming above the bottom in the water column can often avoid toxicity from toxic petroleum compounds in the water column, but if fish are expiring in a confined** habitat (small pond, etc.), EPA GC/MS method 8240 and the NOAA protocol expanded scan*** for PAHs could be performed to see if Benzene, Toluene, Ethyl Benzene, and Xylene (BTEX), naphthalene, and other potentially toxic compounds are above known acute toxicity benchmark concentrations. Zooplankton populations impacted by oil usually recover fairly quickly unless they are impacted in very confined** or shallow environments [835] and the above BTEX and PAH water methods are often recommended rather than direct analyses of zooplankton tissues.
- 11a. The concern is for benthic invertebrates: analyze invertebrate whole-body tissue samples and surrounding sediment samples for polycyclic aromatic hydrocarbons (PAHs) and alkyl PAHs using the NOAA protocol expanded scan***. If the spill is fresh* or the source continuous, risk assessment needs may also require that the sediments which form the habitat for benthic invertebrates be analyzed for Benzene, Toluene, Ethyl Benzene, and Xylene (BTEX) and other volatile compounds using EPA GC/MS

method 8240 or modified EPA method 8240 in the Selected Ion Mode (SIM). Bivalve invertebrates such as clams and mussels do not break down PAHs as well or as quickly as do fish or many wildlife species. They are also less mobile. Thus, bivalve tissues are more often directly analyzed for PAH residues than are the tissues of fish or wildlife.

11b. The concern is for plants or for vertebrate wildlife including birds, mammals, reptiles, and amphibians: polycyclic aromatic hydrocarbons (PAHs) and other petroleum hydrocarbons break down fairly rapidly in many wildlife groups and tissues are not usually analyzed directly. Instead direct effects are investigated and water, soil, sediment, and food items encountered by wildlife are usually analyzed for PAHs and alkyl PAHs using the NOAA protocol expanded scan***. spill is fresh* or the source continuous, risk assessment needs may also require that these habitat media also be analyzed for Benzene, Toluene, Ethyl Benzene, and Xylene (BTEX) and other volatile compounds using EPA GC/MS method 8240 or modified EPA method 8240 in the Selected Ion Mode (SIM). Less is known about plant effects. However, the same methods recommended above for the analyses of water (Step 4 above) and for sediments or soils (Step 6 above) are usually also recommended for these same media in plant or wildlife habitats. If wildlife or plants are covered with oil, see also Step 8 (above) regarding oiling issues.

* Discussion of the significance of the word "fresh": The word "fresh" cannot be universally defined because oil breaks down faster in some environments than in others. In a hot, windy, sunny, oil-microbe-rich, environment in the tropics, some of the lighter and more volatile compounds (such as the Benzene, Toluene, Ethyl Benzene, and Xylene compounds) would be expected to disappear faster by evaporation into the environment and by biodegradation than in a cold, no-wind, cloudy, oil-microbe-poor environment in the arctic. In certain habitats, BTEX and other relatively water soluble compounds will tend to move to groundwater and/or subsurface soils (where degradation rates are typically slower than in a sunny well aerated surface environment). Thus, the judgement about whether or not oil contamination would be considered "fresh" is a professional judgement based on a continuum of possible scenarios. The closer in time to the original spill of nondegraded petroleum product, the greater degree the source is continuous rather than the result of a one-time event, and the more factors are present which would retard oil evaporation or breakdown (cold, no-wind, cloudy, oil-microbe-poor conditions, etc.) the more likely it would be that in the professional judgement experts the oil would be considered "fresh." In other words, the degree of freshness is a continuum which depends on the specific product spilled and the specific habitat impacted. Except for groundwater resources (where the breakdown can be much slower), the fresher the middle distillate oil contamination is, the more one has to be concerned about potential impacts of BTEX compounds, and other

lighter and more volatile petroleum compounds.

To assist the reader in making decisions based on the continuum of possible degrees of freshness, the following generalizations are provided: Some of the lightest middle distillates (such as Jet Fuels, Diesel, No. 2 Fuel Oil) are moderately volatile and soluble and up to two-thirds of the spill amount could disappear from surface waters after a few days [771,835]. Even heavier petroleum substances, such as medium oils and most crude oils will evaporate about one third of the product spilled within 24 hours [771]. Typically the volatile fractions disappear mostly by evaporating into the atmosphere. However, in some cases, certain water soluble fractions of oil including Benzene, Toluene, Ethyl Benzene, and Xylene (BTEX) compounds move down into groundwater. BTEX compounds are included in the more volatile and water soluble fractions, and BTEX compounds as well as the lighter alkanes are broken down more quickly by microbes than heavier semi-volatiles such as alkyl PAHs and some of the heavier and more complex aliphatic compounds. Thus after a week, or in some cases, after a few days, there is less reason to analyze surface waters for BTEX or other volatile compounds, and such analyses should be reserved more for potentially contaminated groundwaters. In the same manner, as the product ages, there is typically less reason to analyze for alkanes using GC/FID techniques or TPH using EPA 418.1 methods, and more reason to analyze for the more persistent alkyl PAHs using the NOAA protocol expanded scan***.

** Discussion of the significance of the word "confined": Like the word "fresh" the word "confined" is difficult to define precisely as there is a continuum of various degrees to which a habitat would be considered "confined" versus "open." However, if one is concerned about the well-being of ecological resources such as fish which spend most of their time swimming freely above the bottom, it makes more sense to spend a smaller proportion of analytical funding for water column and surface water analyses of Benzene, Toluene, Ethyl Benzene, and Xylene (BTEX) and other volatile or acutely toxic compounds if the spill is in open and/or deep waters rather than shallow or "confined" waters. This is because much of the oil tends to stay with a surface slick or becomes tied up in The petroleum compounds which do pass subsurface tar balls. through the water column often tend to do so in small concentrations and/or for short periods of time, and fish and other pelagic or generally mobile species can often swim away to avoid impacts from spilled oil in "open waters." Thus in many large oil spills in open or deep waters, it has often been difficult or impossible to attribute significant impacts to fish or other pelagic or strong swimming mobile species in open waters. Lethality has most often been associated with heavy exposure of juvenile fish to large amounts of oil products moving rapidly into shallow or confined waters [835]. Different fish species vary in their sensitivity to oil [835]. However, the bottom line is that in past ecological assessments of spills, often too much money has been spent on water column analyses in open water settings, when the majority of significant impacts tended to be concentrated in other habitats, such as benthic, shoreline, and surface microlayer habitats.

*** The expanded scan protocols for polycyclic aromatic hydrocarbons (PAHs) and alkyl PAHs have been published by NOAA [828].

End of Decision Tree Key.

It is important to understand that contaminants data from different labs, different states, and different agencies, collected by different people, are often not very comparable (see also, discussion in the disclaimer section at the top of this entry).

As of 1997, the problem of lack of data comparability (not only for water methods but also for soil, sediment, and tissue methods) between different "standard methods" recommended by different agencies seemed to be getting worse, if anything, rather than better. The trend in quality assurance seemed to be for various agencies, including the EPA and others, to insist on quality assurance plans for each project. In addition to quality control steps (blanks, duplicates, spikes, etc.), these quality assurance plans call for a step of insuring data comparability [1015,1017]. However, the data comparability step is often not given sufficient consideration. The tendency of agency guidance (such as EPA SW-846 methods and some other new EPA methods for bioconcentratable substances) to allow more and more flexibility to select options at various points along the way, makes it harder in insure data comparability or method validity. Even volunteer monitoring programs are now strongly encouraged to develop and use quality assurance project plans [1015,1017].

At minimum, before using contaminants data from diverse sources, one should determine that field collection methods, detection limits, and lab quality control techniques were acceptable and comparable. The goal is that the analysis in the concentration range of the comparison benchmark concentration should be very precise and accurate.

It should be kept in mind that quality control field and lab blanks and duplicates will not help in the data quality assurance goal as well as intended if one is using a method prone to false negatives. Methods may be prone to false negatives due to the use of detection limits that are too high, the loss of contaminants through inappropriate handling, or the use of inappropriate methods. The use of inappropriate methods is particularly common related to oil products.

Additional details:

The relative proportions of hazardous compound constituents present in petroleum-based oil contamination is typically quite variable. The farther one progresses from lighter towards heavier oils (the general progression from light towards heavy is the following: Diesel, No. 2 Fuel Oil, Light Crudes, Medium Crude Oils, Heavy Crudes, No. 6 Fuel Oil, etc.) the greater the percentage of PAHs and other semi-volatiles (many of which are not so immediately

toxic as the volatiles but which can result in long term/chronic impacts). Heavier oils such as fuel oil 5 thus need to be analyzed for the semi volatile compounds which typically pose the greatest long term risk, PAHs and alkylated PAHs.

Screening scans: Certain screening scans may be used to monitor the position and magnitude of contamination. Below are a few notes related to screening scans versus distillate fuels:

GC/FID:

While a screening analysis such as GC/FID should be adequate for mid-range products such as diesels, fuel oil no. 2, and possibly jet fuels, lighter gasoline fractions will be lost in a GC/FID analysis (which uses extraction and burning) [657]. Distillate fuels in the C9 to C16 range normally have a boiling range well above the boiling-point of benzene; accordingly, the benzene content of this fraction is usually low [747].

Method 8015:

EPA Method 8015 (for Non-halogenated Volatile Organics) is a gas chromatographic method sometimes recommended for the analysis of volatile and semivolatile compounds. can be used to characterize light and midrange petroleum distillates such as gasoline, diesel, fuel oil, and kerosene. This method can be used to obtain some gross fingerprint information for differentiation between petroleum products, as well as detailed information that can be used to differentiate between different batches of the same product. The major limitation of Method 8015 is its inability to detect nonvolatile compounds. The State of California recommends a "modified method (different from EPA's method 8015 and also different from EPA method 418.1) for gasoline, kerosene, diesel oil, or other fuels in soil and groundwater, as specified in the Leaking Underground Fuel Tank (LUFT) Manual [465].

The California LUFT methods call for packed GC columns which have poor resolving power and make it difficult to obtain detailed information about the hydrocarbon type [810]. Superior GC columns and superior methods (such as ASTM 2887) are available [810]. For example, narrow-bore capillary columns can analzye most of the gasoline, entire diesel fractions, and a substantial portion of the crude oil range [810].

Using the California LUFT manual methods, only an experienced analyst will be able to differentiate diesel fractions from aged gasoline [810]. The oversimplified California methods and models are plagued with many problems [808,810]. Choosing an appropriate solvent for semivolatile analyses always presents a problem; some solvents extract certain compounds better than others and

many present environmental or health risks [810].

HPLC screening scans:

In cases where a less expensive screening scan is desired, consider using an HPLC/Fluorescence scan method for sediment or bile metabolite samples. Such scans are available from laboratories at Texas A. and M., Arthur D. Little, and the NOAA lab in Seattle. This scan is less prone to false negatives and various other problems than some of the more common screening methods (TPH-EPA 418.1 and Oil and Grease). HPLC/Fluorescence is less expensive of more rigorous scans. t.han some the HPLC/fluorescence scan can be used for analyses of fish bile: the scan looks at bile directly for the presence of metabolites of PAHs: naphthalene, phenanthrene, and benzo(a)pyrene. The technique does not identify or quantify actual PAH compounds, but subsequent gas chromatography analyses can be done to confirm the initial findings. Even the semi-quantitative Total Scanning Fluorescence (TSF) done inexpensively by labs such as GERG are a better measure of PAH contamination than GC/FID, which measures less persistent and less hazardous aliphatics.

Additional Pros: HPLC Fluorescence screening methods have been performed extensively by NOAA to locate hotspots for crude oil contamination. NOAA's experience with the Exxon Valdez spill that indicated concentrations of aromatic hvdrocarbons measured bу HPLC/Fluorescence screening were highly correlated with the sums of Aromatic hydrocarbons determined by GC/MS, thus validating the screening method as an effective tool for estimating concentrations of petroleumrelated aromatic hydrocarbons in sediments. Moreover, differences in HPLC chromatographic patterns among sediments suggested different sources of contamination, e.g., crude oil or diesel fuel. Allows crude determinations related to sources: HPLC/Fluorescence analyses allowed at least rough differentiation between aromatic hydrocarbons which may have originated from diesel fuel versus those from boat traffic [521] and The successfully applied procedure was fingerprinting' gasolines, kerosines, diesel oils, heavy fuel oils, lubricating oils, and ship bilge oils [AUTHOR: Saner WA; Fitzgerald GE, PUBLICATION YEAR: 1976 TITLE: Thin-Layer Chromatographic Technique for Identification of Waterborne Petroleum Oils JOURNAL: Environmental Science and Technology SOURCE: Vol. 10, No. 9, p 893-897, September 1976. 6 fig, 4 tab, 7 ref.].

Although EPA method 418.1: Petroleum Hydrocarbons expressed as Total Petroleum Hydrocarbons (TPH), is recommended by many State agencies, some consulting firms, and some laboratories for certain regulatory screening applications (often underground storage tanks), this method is not well suited to fuel oil no. 6 contamination or to the more persistent hazardous constituents in oil. values tend to give the mistaken impression that a site is clean when it really isn't (prone to false For example, a field negatives). test bioremediation of soils contaminated with Bunker C at a refinery in Beaumont, Texas, utilized oil and grease data, which (although the data was quite variable) seemed to indicate bioremediation was taking place [728]. A comparison of the oil and grease data at this site with TPH data at this site suggested the same thing, that the data was quite variable but if anything, the oil was slowly being cleaned up by bioremediation (Bruce Herbert, Texas A. and M., Department of Geology, personal communication, 1995). However, a later study of the same site utilizing the expanded scan for PAHs (a modified EPA 8270 including alkyl homologues and lower detection limits), indicated that very little bioremediation of hazardous alkyl PAHs and multiring PAHs was actually taking place [727]. utilizing either oil and grease or TPH analyses would tend to lead one to the faulty conclusion that the harmful compounds were being naturally cleaned up at an acceptable rate. This is partly because the TPH and oil and grease methods tend to favor the lighter and less alkylated PAHs, whereas many of the carcinogenic and longer lasting PAHs the heavier multi-ringed and alkylated are For more information, see Petroleum compounds. Hydrocarbons entry.

See also: Laboratory and/or Field Analyses section in Oil Spills entry for information on biological indicators of oil exposure. See also: PAHs as a Group entry.